Joint Control of Net Primary Productivity by Climate and Soil Nitrogen in the Forests of Eastern China

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Abstract: The nature and extent of climate and soil nutrient controls in Chinese forests remain poorly resolved. Here, we synthesized the data on carbon–climate–soil in eastern China, and litter N was firstly taken into consideration, to examine the variation of net primary productivity (NPP) and its driving forces. Results showed that NPP had significant latitude pattern and varied substantially across climate zones. Bivariate analyses indicated that mean annual temperature (MAT), mean annual precipitation (MAP), soil N content ($N_{soil}$), and annual litter N ($N_{re}$) were the main controlling factors in spatial pattern of forest NPP. Notably, partial general linear model analysis revealed that MAT, MAP, and $N_{re}$ jointly explained 84.8% of the spatial variation of NPP. Among the three major factors, $N_{re}$ explained more variation of forest NPP than the other two factors, and MAT and MAP affected NPP mainly through the change of litter N rather than via themselves, highlighting the importance of litter N in estimating forest NPP. However, to accurately describe the pattern of forest NPP in China, more detailed field measurements and methodologies on NPP and relevant confounding factors should be addressed in future studies.

Keywords: net primary productivity; climate zone; climate; soil N; litter N

1. Introduction

Net primary productivity (NPP) is a key ecosystem variable and a critical component of the regional and global carbon cycle [1,2]. However, the spatial distribution of forest NPP is still uncertain, due to complicated impacts from various environmental and biological factors, e.g., vegetation distribution, climatic variables, and land use change [3–5]. Therefore, accurate estimation of NPP and its driving forces is essential to understanding terrestrial carbon pools and responses of forest functions to future climate change [6,7].

At a regional scale, NPP is strongly correlated with climatic factors [8–11]. Based on global NPP data, Lieth (1975) [8] developed the climate-driven theory, and described the relationship between climatic factors (annual mean temperature, annual precipitation, and annual evapotranspiration) and NPP in logistic functions. However, it is still unclear whether regional NPP across biomes follows the same pattern [12,13], since the driving factors vary among regions [14–16]. Further, an international coordination for compilation of global NPP data for model validation and development, the Global Primary Productivity Data Initiative (GPPDI), has worked successfully since 1995. However, inadequate observational NPP data seriously inhibit the estimation and modeling of the global carbon cycle and the validation and evaluation of the global carbon models [17].
Globally, forests represent 80% of plant biomass, and 50–60% of annual NPP in terrestrial ecosystems [5,18]. Chinese forests, which cover about half the total land area of China, contain perhaps the widest range of forest types in the world, ranging from boreal forest and mixed coniferous broad-leaved forest in the north, to subtropical evergreen broad-leaved forest, warm temperate coniferous forest, tropical rainforest, and seasonal forest in the south [19,20]. It is regarded that these forests have a significant influence on carbon budget both regionally and globally [21]. A lot of field measurements of forest biomass and NPP estimations are available from multiple sites for the past two decades. However, these data were mostly published in Chinese journals and reports and not accessible to Western scientists. Also, site-based data of forest NPP in China has not yet been synthesized in a consistent manner.

Located in heavily forested area, the North–South Transect of Eastern China (NSTEC) features a high variation in plant composition, climate, and soil substrate materials [22]. It thus provides wide biome heterogeneity to examine spatial pattern of forest NPP. We synthesized the data in the primary literature on NPP in forests within the NSTEC to produce a consistent dataset on NPP, and amassed the data on mean annual temperature (MAT), mean annual precipitation (MAP), soil N content (N_{soil}), and annual litter N (N_{re}). Hobbie (2015) [23] proposed the plant litter feedback paradigm that changes in plant litter traits reinforce patterns of soil fertility and NPP. Thus, we first introduced litter N as available N for plants, which was an improvement over the past studies. Based on these data, we aim to investigate whether temperature and precipitation characterize the pattern of NPP within the transect, to explore whether soil N is a limiting factor for large-scale distribution of NPP, and to test the applicability of the climate-driven theory in eastern China. Besides understanding the causes of variability in ecosystem productivity, the findings are crucial for assessing the potential responses to global climatic change, and are thus incorporated into statistical and simulation models.

2. Materials and Methods

2.1. Study Area

NSTEC provides an ideal platform for exploring the growth of forest in East Asia’s monsoon region [22]. NSTEC has a spatial distance of more than 3700.0 km in length, ranging from 108.0° E to 118.0° E for latitude below 40.0° N and from 118.0° E to 128.0° E for latitude above 40.0° N (Figure 1). From north to south, MAP increases from 500.0 mm to about 1800.0 mm, and MAT changes from 1.0 to 22.0 °C, correspondingly. Due to the obvious latitudinal gradients for climate, zonal forest ecosystems occur within the NSTEC, which include cold-temperate coniferous forest, temperate mixed forest, warm-temperate deciduous broadleaved forest, subtropical evergreen broad-leaved forest, and tropical monsoon rainforest. Within NSTEC, 87 observations in 34 plots were included in this study, and geographical distribution of the sites was mapped in Figure 1.

2.2. Datasets

Data on the forest ecosystems within the transect were obtained from published literature (Figure 1, Table S1). The sampling years were not mentioned in most of the original literature, and only the publication years of the literature could be obtained, which varied from 1983 to 2010. The information about the publication time could be seen in Table S1. The times were classified into three groups according to the publication year, and they were 1980s (1983–1989), 1990s (1991–1999), and 2000s (2001–2010). Based on the classification, the temporal variations were analyzed, and there were no significances among climatic factor, soil N, and also NPP for different times (Table S2). Therefore, we take no account of temporal variations of forest NPP, and only spatial pattern of forest NPP was involved. To be clear, “major forest biome” was used to represent a higher-level classification of the “ecological zone”, which included boreal, temperate, subtropical, and tropical forests [24]. Each site included site name, latitude, longitude, tree species, MAT, MAP, N_{soil}, N_{re}, and NPP estimations (Table S1). We retrieved missing latitude or longitude information for sites without such data from
Google Earth according to site names. Meanwhile, missing MAT and MAP were extracted from the Chinese climate data based on site locations. NPP was expressed, herein, in per unit of oven-dry matter (t ha\(^{-1}\) a\(^{-1}\)). In order to compare with other studies, NPP may be expressed in g C m\(^{-2}\) a\(^{-1}\), where 1.0 g carbon is equivalent to 2.2 g oven-dry matter.

![Figure 1](image-url) Locations of research plots in this synthesis in eastern China (black dots). The region between the two lines represents the area of the North–South Transect of Eastern China (NSTEC).

### 2.2.1. Estimation of NPP

Forest NPP was estimated as the sum of increase in the standing crop of vegetation based on the data for biomass, which included net increments of trees, shrubs, and herbs. Biomass data in our study were all from field survey, and data from modeling and regional average were excluded. The sampling methods for measuring forest biomass used by Chinese scientists, however, were quite different. General methods of biomass estimates were given below. Firstly, diameter at breast height (DBH) and the height (H) of each tree in the plot were measured, and then one of the three methods was used: (1) mean tree biomass was measured, and then multiplied by tree density of each plot; (2) allometric equations were used to calculate the biomass of each tree; and (3) allometric equations were used to calculate the mean tree biomass, and multiplied by tree density of each plot [25,26]. Secondly, dead biomass of each tree was the sum of standing dead stem and coarse woody debris [27]. Thirdly, biomass of shrubs and herbs was measured by harvest method within large tree samples. Aboveground tissues were clipped, and the weight of these tissues and the aboveground biomass were calculated based on the plot area. Finally, the biomass of litterfall was determined by monthly collection in three or more 1.0 m × 1.0 m plots, laid out inside a single tree plot. Litterfall components, such as leaf, branch, flower, and fruit, were dried and weighed to estimated total litterfall biomass. After one year, the sampling and the analysis for trees and understory vegetation were repeated, and the increases in biomass were taken as forest NPP.

### 2.2.2. Measurements of Soil N

Soil sampling depths varied from a few centimeters to several meters in different studies. Considering data availability and distribution of plant roots in the soil, in our study, only the data from the depths of 0–60.0 cm in soil were used. Generally, the soil was sampled from the soil pits selected randomly in each forest ecosystem, and soil bulk density was determined by collecting samples in volumetric rings. In preparation for analysis, soil samples were air-dried and then passed through the mesh sieve for measurement of N concentration. \( N_{\text{soil}} \) was calculated by multiplying the mean concentration of N in each layer with the corresponding mean soil bulk density Equation (1).
Annual litter N (\(N_{re}\)) was calculated as litter biomass multiplied by the respective N concentration, and finally, the contents of the individual fractions were summed in Equation (2).

\[
N_{soil} = \sum (\text{mean concentration of N in each layer} \times \text{corresponding mean soil bulk density}) \quad (1)
\]

\[
N_{re} = \sum (\text{annual litter biomass of each component} \times \text{litter N concentration in each component}) \quad (2)
\]

where, \(N_{soil}\) is soil N content (kg ha\(^{-1}\)), \(N_{re}\) is annual litter N (kg N ha\(^{-1}\) a\(^{-1}\)).

2.2.3. Climate Data

When investigating the relationship between climate and forest productivity, it would be better to employ climate data for particular sampling years. Hence, synchronous climate data for particular sampling years were collected. However, most sampling sites were far away from weather stations, and thus, climate data were missing. Therefore, the meteorological data from the Meteorological Database of the Chinese Ecological Research Network (CERN) Synthesis Research Center were extracted. Climate data in a grid cell where the sampling site was located was extracted as the climate data of the sampling site. MAP and MAT were the average values of 1980–2000 with a 10-day-0.1° spatial–temporal resolution [28].

2.3. Statistical Analysis

One-way analysis of variance (ANOVA) was adopted to test the differences in forest NPP among different climate zones, and was followed by Fisher’s least significant difference (LSD) comparisons when the differences were significant. Linear regression analysis or dynamic curve fit was used to analyze the relationships between forest NPP and the driving forces. Considering the differences in sample size, we compared \(R^2\) and root mean squared error (RMSE), and selected the better-fit functions that had a higher \(R^2\) and lower RMSE. The stepwise regression was used to analyze the linear regression on forest NPP with climate and soil N. In the stepwise regression, the minimum \(p\)-value for a variable to be recommended for adding to and removing from the model was 0.05. Considering the results of stepwise regression, only MAT, MAP, and \(N_{re}\) were analyzed in the ensuing analysis. To identify the relative effects and interactive effects of the above three factors on forest NPP, we conducted a partial general linear model (GLM) using NPP as dependent variable, and MAT, MAP, and \(N_{re}\) as predictors. The partial regression divides the variation in response variable explained by several predictor variables into independent components (representing the independent effects of an individual explanatory variable when controlling effects of the other explanatory variables) and joint components (usually representing the collinearities between explanatory variables). The variation partitioning with three explanatory matrices leads to the identification of seven fractions in this study, i.e., independent effects of MAT, MAP, and \(N_{re}\); interactive effects of MAT and MAP, MAT and \(N_{re}\), MAP and \(N_{re}\), and the interactive effect of all variables. Further details about the method were given in Heikkinen et al. (2005) [29].

Figure 1 was plotted with ArcGIS 10.1 software (Esri, Re lands, CA, USA), and other graphs were performed by Sigma Plot 13.0 software (Systat Software Inc., San Jose, CA, USA). The partial GLM was performed with SAS statistical software and other analyses were conducted by SPSS 16.0 statistical software (SPSS Inc., Chicago, IL, USA).

3. Results

3.1. Latitudinal Pattern and Statistics of NPP

Forest NPP across eastern China exhibited significantly obvious latitudinal patterns \((p < 0.001)\), but no clear longitudinal trend \((p = 0.20)\) (Figure 2). Therefore, we did not elaborate on the longitudinal pattern of forest NPP. Generally, NPP decreased with increasing latitude when latitude was below 35.0° N, but increased when the latitude was higher than the threshold level (Figure 2). As a whole,
mean total NPP was 9.5 ± 0.7 t ha\(^{-1}\) a\(^{-1}\) (mean ± SE), ranging from 0.8 to 29.6 t ha\(^{-1}\) a\(^{-1}\), with the variability (coefficient of variation, CV) up to 64.0% (Table 1). In terms of different climate zones, tropical forests below 23.0° N had the highest forest productivity with an average NPP of 15.3 ± 1.3 t ha\(^{-1}\) a\(^{-1}\), but the smallest spatial variability (CV = 31.6%). NPP of temperate forests was, on average, 5.5 ± 0.8 t ha\(^{-1}\) a\(^{-1}\), which tended to be the lowest among the four climate zones. The other climate zones (subtropical forests and boreal forests) were not different from each other, and were intermediate to the others. Specifically, the average NPP of subtropical forests was 10.1 ± 0.9 t ha\(^{-1}\) a\(^{-1}\), which was close to the value of boreal forests (12.1 ± 4.7 t ha\(^{-1}\) a\(^{-1}\)), but the latter forests exhibited a higher spatial variability in NPP from available data (CV = 78.4%) (Table 1).

![Figure 2](image_url)

**Figure 2.** Trends of net primary productivity (NPP) along latitude for forest ecosystems in eastern China. N, number of observations; F, F Values.

### Table 1. Comparisons of net primary productivity (NPP) of forest ecosystems in different climate zones.

<table>
<thead>
<tr>
<th>Climate Zone</th>
<th>N</th>
<th>Mean (t ha(^{-1}) a(^{-1}))</th>
<th>Max (t ha(^{-1}) a(^{-1}))</th>
<th>Min (t ha(^{-1}) a(^{-1}))</th>
<th>SE</th>
<th>CV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boreal</td>
<td>4</td>
<td>12.1 ab</td>
<td>24.6</td>
<td>1.9</td>
<td>4.7</td>
<td>78.4</td>
</tr>
<tr>
<td>Temperate</td>
<td>28</td>
<td>5.5 c</td>
<td>14.1</td>
<td>0.8</td>
<td>0.8</td>
<td>73.2</td>
</tr>
<tr>
<td>Subtropical</td>
<td>42</td>
<td>10.1 b</td>
<td>29.6</td>
<td>1.2</td>
<td>0.9</td>
<td>54.7</td>
</tr>
<tr>
<td>Tropical</td>
<td>13</td>
<td>15.3 a</td>
<td>24.2</td>
<td>8.6</td>
<td>1.3</td>
<td>31.6</td>
</tr>
<tr>
<td>Overall</td>
<td>87</td>
<td>9.5</td>
<td>29.6</td>
<td>0.8</td>
<td>0.7</td>
<td>64.0</td>
</tr>
</tbody>
</table>

3.2. **The Impact of Climatic Factors on the Spatial Pattern of NPP**

Temperature is an important driving factor for the ecosystem carbon budget, and water is the basic material for maintaining ecosystem structure and functions. As was clearly shown in Figure 3, the plot based forest NPP grew linearly with increasing MAT, MAP, and the combinations, although variation occurred within the climate band and few sites fell out of 95.0% prediction band. On average, MAT and
MAP contributed 16.6% and 21.8% of the spatial variation of NPP, respectively (Figure 3, Table 2). We then analyzed the combined contribution of MAT and MAP to the spatial variation of NPP, and found that MAT and MAP jointly explained 24.3% of the spatial variation of NPP, which was only just 2.5% higher in the prediction of NPP than single climatic factor (MAP) alone (Figure 3, Table 2). As for different climate zones, climatic factors were also closely related to the occurrence of NPP. Generally, there was a consistent and significant shift from species with high NPP in warm climate with high MAT and MAP toward species with low NPP in low MAT and MAP conditions, while the trend in boreal forest ecosystems differed (Figure S1). Specifically, for boreal forest, higher NPP occurred with lower MAT and MAP. More speculatively, soil nutrients or other environmental factors might contribute to the variation of forest NPP in this region, and this should be taken into consideration in the ensuing analyses. It is worth noting that the number of boreal forests was quite low, hence, the reliability of the results should be considered carefully.

![Figure 3. Relationships between net primary productivity (NPP) and mean annual temperature (MAT), mean annual precipitation (MAP), and combinations of MAT and MAP. N, number of observations; F, F Values.](image)

Table 2. Bivariate and multivariate regression models of net primary productivity (NPP) on climatic factors and soil N. MAT, mean annual precipitation; MAP, mean annual temperature; N$_{soil}$, soil N content; N$_{re}$, annual litter N; N, number of observations; F, F Values; RMSE, root mean square error.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Model Type</th>
<th>N</th>
<th>$R^2$</th>
<th>F</th>
<th>RMSE</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAT</td>
<td>linear</td>
<td>87</td>
<td>0.166</td>
<td>16.9</td>
<td>5.6</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>exponential</td>
<td>87</td>
<td>0.202</td>
<td>22.0</td>
<td>5.6</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>MAP</td>
<td>linear</td>
<td>87</td>
<td>0.218</td>
<td>21.5</td>
<td>5.4</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>exponential</td>
<td>87</td>
<td>0.237</td>
<td>23.2</td>
<td>5.3</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>MAT+MAP</td>
<td>linear</td>
<td>87</td>
<td>0.243</td>
<td>23.8</td>
<td>4.2</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>exponential</td>
<td>42</td>
<td>0.106</td>
<td>4.7</td>
<td>7.0</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>N$_{soil}$</td>
<td>linear</td>
<td>42</td>
<td>0.148</td>
<td>4.5</td>
<td>6.7</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td></td>
<td>exponential</td>
<td>86</td>
<td>0.284</td>
<td>30.1</td>
<td>3.4</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>N$_{reg}$</td>
<td>linear</td>
<td>42</td>
<td>0.382</td>
<td>32.7</td>
<td>2.2</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>N$<em>{soil} + N</em>{reg}$</td>
<td>linear</td>
<td>42</td>
<td>0.543</td>
<td>10.6</td>
<td>1.6</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>MAT + MAP + N$<em>{soil} + N</em>{reg}$</td>
<td>linear</td>
<td>42</td>
<td>0.543</td>
<td>10.6</td>
<td>1.6</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

3.3. The Impact of Soil N on the Spatial Pattern of NPP

Similar to climatic factors, our data illustrated that soil N was also closely related to forest NPP (Figure 4). Generally, positive linear function fitted well the relationship between forest NPP and soil
N, and namely, N_{soil} and N_{re} accounted for 10.6% and 28.4% of variation of NPP (Figure 4, Table 2). When examining the combined effect of N_{soil} and N_{re}, we found that N_{soil} and N_{re} in the model jointly explained 38.2% of variation in NPP, a marked improvement over N_{soil} or N_{re} alone. (Figure 4, Table 2). Although forest NPP covaried spatially with soil N, it should be noted that a statistically significant relationship did not necessarily imply causality. Moreover, the occurrence of high N_{soil} and N_{re} rather than MAT and MAP favored the increase of NPP in boreal forest ecosystems (Figure S2). Additionally, across all bivariate analyses above, $R^2$-value and $F$-value of N_{re} for NPP were almost always higher than those of NPP versus other factors involved in this study, illustrating that plant litter N played a more important role in determining forest NPP than other environmental factors.

![Figure 4](image.png)

**Figure 4.** Relationships between net primary productivity (NPP) and soil N content ($N_{soil}$), annual litter N ($N_{re}$), and combinations of $N_{soil}$ and $N_{re}$. $N$, number of observations; $F$, $F$ Values.

### 3.4. The Combined Impact of Climate and Soil on the Spatial Pattern of NPP

Even though the climate-NPP and soil-NPP relationships analyzed above were statistically significant, many points were still scattered around the fitted lines, and a great deal of variability for NPP was not captured (Figures 3 and 4). Subsequently, we conducted a stepwise multiple regression to identify the effects of climatic factors and soil N on forest NPP. The results showed that $N_{soil}$ was excluded from the linear regression model, and the explanatory power of these three factors (MAT, MAP, and $N_{re}$) for NPP was just 49.2%, a marginal decline over four factors together, in Equation (3).

$$NPP = -0.3387 - 0.033 \text{MAT} + 0.004 \text{MAP} + 0.073N_{re}$$

$$R^2 = 0.492, N = 42, F = 10.2, p < 0.001$$  \hspace{1cm} (3)

However, there were significant collinearities between these environmental factors. Table 3 summarized correlation coefficients among these variables. Climatic factors for MAT and MAP were highly correlated, and both of them had marked correlations with $N_{re}$, but had non-robust correlation with $N_{soil}$, and mutually, $N_{re}$ had a marginal relationship with $N_{soil}$ (Table 3).
Considering the results of stepwise multiple regression analysis and correlation coefficients above, only MAT, MAP, and N<sub>re</sub> were referred to when identifying the combined effects of the drivers on NPP in ensuing analysis. Given the significant collinearities among the three factors, their true roles for NPP could be obscured. Therefore, we used partial GLM to examine their relative causality in the control of NPP in forests. General linear model involving MAT, MAP, and N<sub>re</sub> could account for 84.8% of the variation in NPP, and MAT, MAP, and N<sub>re</sub> explained 9.8%, 12.7%, and 35.3% of the variation in NPP, respectively; the interactive effects of MAT and MAP (ab), MAT and N<sub>re</sub> (ac), MAP and N<sub>re</sub> (bc), and MAT, MAP, and N<sub>re</sub> (abc) represented 10.9%, 4.6%, 4.0%, and 7.5%, respectively (Figure 5). The results indicated that N<sub>re</sub> was much more important in shaping forest NPP than the climatic factors. Full model for NPP:

\[
\text{NPP} = a \text{MAT} + b \text{MAP} + c \text{N}_{\text{re}} + \text{abc} \text{MAT} \times \text{MAP} \times \text{N}_{\text{re}}
\]

\[
\text{Total} = a \text{MAT} + b \text{MAP} + c \text{N}_{\text{re}} + \text{abc} \text{MAT} \times \text{MAP} \times \text{N}_{\text{re}}
\]

4. Discussion

4.1. Carbon Budget

NPP in forests in eastern China in our study were, on average, 4.3 t C ha<sup>-1</sup> a<sup>-1</sup>, which was similar to that in boreal forest ecosystems (4.2 t C ha<sup>-1</sup> a<sup>-1</sup>) [30], but lower than that in European Forest Ecosystems (6.5 t ha<sup>-1</sup> a<sup>-1</sup>) [31]. Actually, considering the number and spatial representation...
of observation sites in different regions, our results need to be further validated with more data. Based on the 7th national forest inventory data and biomass–volume relationship, we estimated that forest biomass carbon in China was approximately 840.3 Tg C a$^{-1}$ (forest area was 195.0 $\times$ 10$^6$ ha), assuming that forests represented potential vegetation in forest regions, and the impact of human disturbance was negligible. Our findings were similar to the result reported by Ni et al. (2003) [20] (738.9 Tg C a$^{-1}$), but was about two times that estimated by Fang et al. (2001) [19] (461.0 Tg C a$^{-1}$), and the results might be ascribed to the differences in NPP estimate methods and forest area calculations. As a large country in the world, China contributes much to regional and global carbon budget. Therefore, determining distribution pattern of carbon budget of the forest ecosystems in China will be helpful for exploring the carbon cycle of the terrestrial ecosystem and addressing global warming [32].

4.2. Explanations for the Distribution Pattern of NPP

4.2.1. Climate Control

Temperature is an important factor for regulating potential photosynthetic activity of vegetation, as well as the length of the growing season [33,34], both of which jointly determine ecosystem productivity [35]. In addition to temperature, precipitation is another determinant of the spatial variation of NPP [36,37]. Numerous studies have indicated that NPP increased with both increasing temperature and precipitation, but the response rates and patterns varied in different regions [9,10,36]. Our study found that NPP in the forest ecosystems of China increased linearly at a rate of 0.38 t ha$^{-1}$ a$^{-1}$ for a 1.0 $^\circ$C increase in MAT, and 1 t ha$^{-1}$ a$^{-1}$ for a 1.0 mm increase in MAP (Figure 3). Our results were similar to that of Ni et al. (2001) [38], who noted that NPP in the forest ecosystems of China increased by 0.48 kg ha$^{-1}$ a$^{-1}$ for 1.0 $^\circ$C increase in MAT and 0.1 kg ha$^{-1}$ a$^{-1}$ for 1.0 mm increase in MAP. Luo et al. (2004) [10] found that NPP increased exponentially, rather than linearly, with an increase in MAT, whereas NPP increased with increasing MAP when MAP was lower than the threshold level of 1490.0 mm, then decreased when MAP was higher than the threshold level. Luyssaert et al. (2007) [36] illustrated that in a global forest study, NPP increased with increasing MAT from 5.0 to 10.0 $^\circ$C, but appeared to be saturated beyond 10.0 $^\circ$C. Similarly, NPP increased with increasing MAP until leveling off at 1500.0 mm. Although the responses of NPP to climatic factors were different in a certain extent, the decisive effects of temperature and precipitation on spatial variation of NPP were established in most studies.

4.2.2. N Control

Soil N is a primary factor that limits plant growth owing to the large discrepancy between demand and supply, and its vital role in plant carbon assimilation [39,40]. Significant advances have been made in the past decades toward understanding the relationship between soil N and NPP in terrestrial ecosystems, whereas substantial uncertainties persist, and discrepancies between studies remain unresolved [40–42]. Yuan et al. (2006) [43] found a significant positive linear correlation between NPP and soil inorganic N for grassland ecosystems in Mongolia, whereas Luo et al. (2004) [10] found a curved relationship between NPP and total soil N. Our field data indicated that N$_{\text{soil}}$ alone explained nearly 10.6% of the NPP variation within the transect (Figure 4, Table 2). Hobbie (2015) [23] focused on feedback to NPP operating through litter decomposition, and reported that positive or negative effects of litter N on the later stages of litter decomposition could strengthen or weaken the positive loop of NPP. Therefore, beside N$_{\text{soil}}$, we introduced N$_{\text{re}}$ to estimate the effect of soil N on forest NPP, and found that N$_{\text{re}}$ alone accounted for 28.4% of the variation, which had higher explanation for NPP than N$_{\text{soil}}$, and the positive effects of N$_{\text{re}}$ on forest NPP were confirmed (Figure 4, Table 2). The role of soil N in shaping forest NPP, in our study, was imperative for models based on mechanism and ecological process, such as CEVSA model, and BIOME-BGC model for NPP estimation. Generally, the plant growth module in CEVSA model described photosynthesis, carbon allocation, leaf area index,
and litter production; the biogeochemical module simulated the transformation and decomposition of organic materials, and nitrogen inputs and outputs in soil. BIOME-BGC model was described as the function of temperature, leaf area, water, and soil nitrogen.

Soil N and litter N were involved in the key process of these models. Thus, future efforts should be focused on identifying the role of soil N, litter N, litter nutrient limiting decomposition, and litter nutrient release versus NPP.

4.2.3. Joint Control

The present study identified one geographical trend in forest productivity, and found that MAT and MAP, alone or in combination, explained 16.6–24.3% of the NPP variation within the transect, while \(N_{\text{soil}}\) and \(N_{\text{re}}\) explained 10.6–38.2% of the variation (Figures 3 and 4, Table 2). Although forest NPP covaried spatially with climatic factors and soil N, it should be noted that a statistically significant relationship did not necessarily imply causality. Importantly, we found that the trends of the relationships of NPP to soil N were more similar than that of NPP versus climatic factors, especially for boreal forests, indicating that rich soil N, rather than warm climate, stimulated the increase in productivity in this region (Figures 3 and 4). To confirm the effects of climate and soil on NPP, partial GLM was conducted, and the results showed that the overall model including MAT, MAP, and \(N_{\text{re}}\), could account for 84.8% of the spatial variation in NPP (Figure 5). The findings suggest that the combination of temperature and precipitation-related physiology and soil substrate-related N is responsible for the observed pattern of forest productivity, and also reflects pervasive geographic pattern in the structure and function of forest ecosystems [42]. It was worth noting that in the overall model, \(N_{\text{re}}\) explained independently more than 35.0% of the total variation for NPP, while MAT and MAP explained independently less than 15.0% (Figure 5). The results above suggest that the geography of NPP was largely controlled by \(N_{\text{re}}\) rather than climate, highlighting the role of litter N in shaping the spatial pattern of forest NPP. The finding is valuable for planners and decision-makers in their attempts to evaluate the effects of nutrient status on forest ecosystems and to develop suitable strategies for forest management.

4.3. Uncertainty Analysis

In this study, we synthesized the data in the primary literature on NPP in the forest ecosystems in eastern China to produce a consistent dataset on NPP. With a wide coverage and a reasonable range, the results of this study provide a robust estimate of NPP in this region. However, there was still a great deal of variability in NPP, which might be ascribed to limitations of data and methodology. Firstly, new methods of data collection are still urgently required. Data collection should come directly from the ecologists who measure forest NPP, rather than indirectly from publications. Standard methods in measuring biomass, standard measure time, and suitable methods in estimating NPP are strongly encouraged. Secondly, uneven site distribution could be an important factor contributing to the uncertainty. In this study, the sites tended to be more concentrated in the temperate and subtropical than boreal zone; therefore, limitations in the results of the analysis were introduced by this uneven site distribution. Additionally, when analyzing the spatial variation of NPP at an annual scale, errors were likely to be produced due to different time spans of data collection. Moreover, available field measurements of NPP depend largely on the sum of the positive increments of biomass, since root production was rarely estimated. As a result, existing field-based estimates of forest NPP in our study were likely to be significant underestimates. Furthermore, it is worth noting that although MAT, MAP, and \(N_{\text{re}}\) were important factors in the spatial variation of NPP in the forests of eastern China, the total variability that was not captured was 15.2% for NPP (Figure 5). We inferred that NPP showed high spatial variability due to impacts from various environmental factors, and biological factors, such as stand age, vegetation distribution, nitrogen deposition, and disturbance (i.e., thinning harvesting irrigation, and drainage) [44–46], could weaken the climate and soil regulation on forest productivity to a certain extent. An intensive study on this aspect is expected to be carried out in the future,
which is critical for expanding current analysis and including other carbon cycles related to more environmental factors.

5. Conclusions

Based on published data from Chinese literature and reports, we concluded that NPP increased from north to south within the NSTEC, and high spatial variation of NPP was found among climate zones. Spatially, climate and soil were both significantly linearly correlated to NPP, and the combined effect of MAT, MAP, and \( N_{\text{re}} \) accounted for 84.8% of the spatial variation of NPP. Considering the true roles of the controlling factors in NPP, \( N_{\text{re}} \) was the most important, followed by MAP and MAT in succession. The findings demonstrate that forest productivity was determined by climate, mainly via the status of soil N, highlighting the role of litter N in shaping the spatial pattern of forest NPP. The results are helpful for elucidating spatial variation of NPP and evaluating potential responses of forest ecosystems to global climatic change. However, further studies need to be carried out to fully understand and verify the causes of variability of NPP in forest ecosystems across wider geographical sites.

Supplementary Materials: The following materials are available online at http://www.mdpi.com/1999-4907/9/6/322/s1. Table S1: Location, mean annual temperature (MAT), mean annual precipitation (MAP), soil N content (\( N_{\text{soil}} \)), annual litter N (\( N_{\text{re}} \)) and net primary productivity (NPP) in this study. NOTES: Table S1 is a long table, and we put it in a separate file; Table S2: Summary of analysis of variance (ANOVA) of mean annual temperature (MAT), mean annual precipitation (MAP), soil N content (\( N_{\text{soil}} \)), annual litter N (\( N_{\text{re}} \)) and net primary productivity (NPP) among three classification of publication year (1980s (1983–1989), 1990s (1991–1999) and 2000s (2001–2010)). Figure S1: Comparisons of mean annual temperature (MAT), mean annual precipitation (MAP), and net primary productivity (NPP) of forest ecosystems in different climate zones. Different letters above bars indicated significant differences among climate zones, which were determined by Fisher’s least significant difference (LSD) comparisons (\( p < 0.05 \)). Figure S2: Comparisons of soil N content (\( N_{\text{soil}} \)), annual litter N (\( N_{\text{re}} \)), and net primary productivity (NPP) of forest ecosystems in different climate zones. Different letters above bars indicated significant differences among climate zones, which were determined by Fisher’s least significant difference (LSD) comparisons (\( p < 0.05 \)).

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